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Citation

Weingärtner, Sebastian, Mehmet Akcakaya, Sophie Berg, Kraig V Kissinger, Warren J Manning, and Reza Nezafat. 2013. Heart-rate independent myocardial t1-mapping using combined saturation and inversion preparation pulses. *Journal of Cardiovascular Magnetic Resonance* 15(Suppl 1): P46.

Published Version

doi:10.1186/1532-429X-15-S1-P46

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POSTER PRESENTATION

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Heart-rate independent myocardial T1-mapping using combined saturation and inversion preparation pulses

Sebastian Weingärtner^{1,3*}, Mehmet Akcakaya¹, Sophie Berg¹, Kraig V Kissinger¹, Warren J Manning^{1,2}, Reza Nezafat¹

From 16th Annual SCMR Scientific Sessions

San Francisco, CA, USA. 31 January - 3 February 2013

Background

Myocardial T1 mapping remains a challenging task due to restrictions imposed by cardiac and respiratory motion. Modified Look-Locker Inversion Recovery (MOLLI) [1] is widely used for 2D cardiac T1-mapping. In MOLLI, the spin-lattice relaxation curve is sampled several times after a single magnetization preparation. The ECG triggered imaging induces a disturbance in the relaxation curve, which varies based on the heart rate. Hence, MOLLI T1 measurements show strong correlations to the heart rate especially in pre-contrast. We developed a novel T1

mapping sequence that enables heart-rate invariant myocardial T1 mapping.

Methods

Figure 1 shows the schematic of the proposed SATuration Pulse Prepared Heart rate independent Inversion-REcovery sequence (SAPPHIRE). A saturation pulse is inserted right after the R-wave of selected heart-cycles. This dephases the magnetization in the imaging volume and eliminates the need for recovery periods after the magnetization preparation. The saturation pulse is followed by an

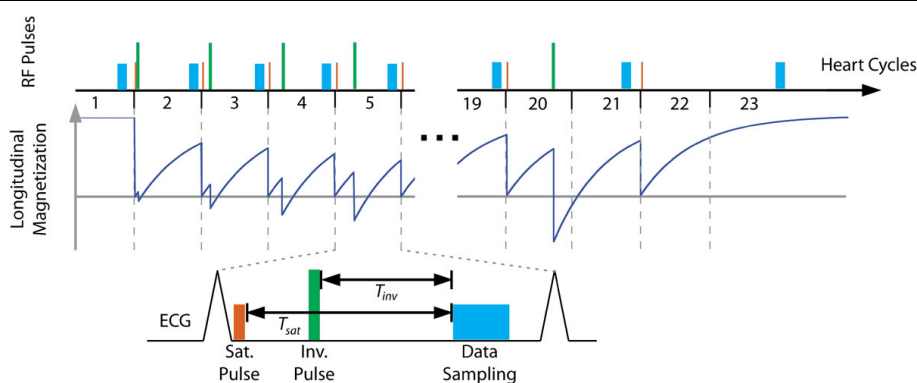


Figure 1 Sequence diagram depicting the SAPPHIRE T1-mapping sequence: a saturation pulse is performed after the R-wave to erase the magnetization history. It is followed by the inversion pulse and a single-shot image readout. To extend the range of applicable inversion times the data readout of some SAPPHIRE experiments is performed in the heart-cycle after the magnetization preparation. Additionally the first heart-cycle is performed without magnetization preparation and the last heart-cycle with the saturation pulse only. This increases the effective inversion times and improves the T1 fit.

¹Department of Medicine, Beth Israel Deaconess Medical Center and Harvard Medical School, Boston, MA, USA

Full list of author information is available at the end of the article

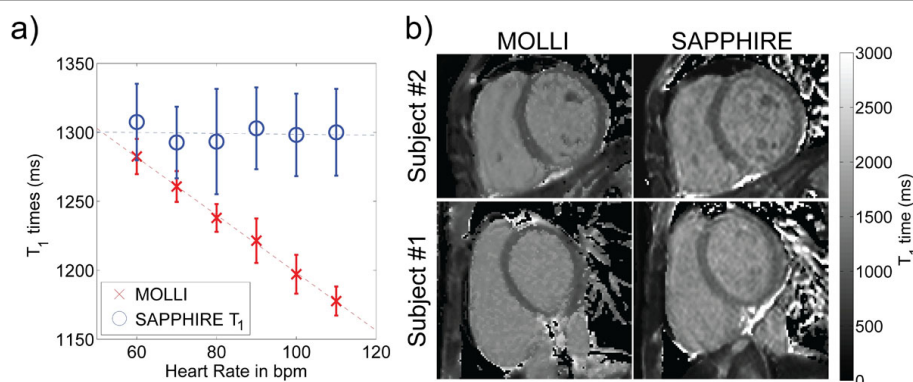


Figure 2 a) Calculated T1 values from phantom images acquired with different simulated R-R interval of 60-120 beats per minute. T1 measurements calculated from MOLLI shows linearly dependent on the duration of R-R, however SAPHIRE T1 measurements were relatively constant. b) T1 maps from two healthy subjects acquired without gadolinium contrast showing higher homogeneity of the T1 measurements in SAPHIRE.

inversion pulse after a variable delay to create various T1 weighted contrasts in the images. Eleven SAPHIRE images are acquired, where each magnetization preparation is followed by a single-shot imaging in the same heart-cycle. Six additional SAPHIRE images are acquired with longer inversion times, by performing the data sampling in the heart-cycle after the magnetization preparation. The first heart cycle is performed without any prepulses, to provide a spin-density weighted image, which facilitates the T1-fit.

SAPHIRE T1-mapping was compared to MOLLI in phantom measurements and in healthy volunteers. A bottle phantom with a T1 of ~1300 ms was imaged using both T1-mapping sequences at various simulated ECGs with different heart-rates. Furthermore, pre-contrast T1-maps in five healthy volunteers were acquired using SAPHIRE T1-mapping and MOLLI.

Results

In the phantom measurements SAPHIRE T1-mapping is in good agreement with MOLLI measurements at a simulated heart-rate of 60 bpm (Relative difference: <2%). The SAPHIRE T1-times, as depicted in Figure 2a), showed no significant correlation with the heart rate ($r = -0.10$), while MOLLI is highly correlated ($r = -0.99$). The T1 times in myocardium and the blood pool of the LV of the volunteers showed no significant difference between the two sequences ($p = 0.20$, $p = 0.10$). Figure 2b) shows exemplary T1-maps of two subjects. SAPHIRE T1-mapping required slightly longer breath holds (16-23s SAPHIRE vs. 12-17s MOLLI).

Conclusions

SAPHIRE T1-mapping enables heart rate independent myocardial T1-mapping. The heart-rate invariance is

achieved by applying a combination of saturation and inversion pulses as magnetization preparation.

Funding

Deutsche Telekom Stiftung; NIH:R01EB008743-01A2; NIH: K99HL111410-01.

Author details

¹Department of Medicine, Beth Israel Deaconess Medical Center and Harvard Medical School, Boston, MA, USA. ²Department of Radiology, Beth Israel Deaconess Medical Center and Harvard Medical School, Boston, MA, USA. ³Computer Assisted Clinical Medicine, University Medical Center Mannheim, Heidelberg University, Mannheim, Germany.

Published: 30 January 2013

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doi:10.1186/1532-429X-15-S1-P46

Cite this article as: Weingärtner et al.: Heart-rate independent myocardial T1-mapping using combined saturation and inversion preparation pulses. *Journal of Cardiovascular Magnetic Resonance* 2013 **15** (Suppl 1):P46.

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